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Reply to Office Action of November 7, 2006

AMENDMENTS TO THE CLAIMS

This listing of claims will replace all prior versions, and listings, of claims in the present

application.

Listing of Claims:

1. (Previously Presented) A composition for improving lipid metabolism having

lactoferrin as an active ingredient.

2. (Previously Presented) A composition for treating at least one disease to be selected

from the group consisting of hypercholesterolemia, hyper-neutral lipidemia, hyper-low density

lipoprotein (LDL) cholesterolemia, hypo-high density lipoprotein (HDL) cholesterolemia,

obesity, fatty liver and cholesterol gallstone which has lactoferrin as an active ingredient.

3. (Previously Presented) A composition for enhancing basal metabolic rate which has

lactoferrin as an active ingredient.

4. (Currently Amended) The composition of any one of claims 1 to 3, wherein said

composition which is in the form of a dusting powder, a powder, a granule, a tablet or a capsule.

and can be obtained by the steps comprising mixing the active ingredient with pharmaceutically

acceptable additives in the dry state; if desired, subjecting the mixture to strong pressure molding in the dry state and successively forming the molded product into fine particulates or granules of

a uniform size; and tableting or encapsulating the mixture, the fine particulates or granules.

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5. (Previously Presented) The composition of claim 1 which is in the form of an enteric

coated preparation.

6. (Previously Presented) The composition of claim 1, wherein tableted granules

containing the active component is coated with a film having, as the major component, a base

which has resistance to the gastric juice and dissolves in the small intestine.

7. (Currently Amended) The composition of claim 1, wherein which is for the amount

of administered administration of the active ingredient is in an amount of about 0.1 mg to about

50,000 mg, preferably about 0.5 mg to about 10,000 mg, more preferably about 10 mg to about

2,000 mg a day.

8. (Withdrawn) A method for producing a composition of claim 1 comprising the steps

of mixing the active ingredient with pharmaceutically acceptable additives in the dry state; if

desired, subjecting the mixture to strong pressure molding in the dry state and successively

forming the molded product into fine particulates or granules of a uniform size; and tableting or

encapsulating the mixture, the fine particulates or granules, said composition being in the form of

a dusting powder, a powder, a granule, a tablet or a capsule.

9. (Currently Amended) Use of A method of improving lipid metabolism, said method

comprising:

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administering the composition of claim 1 having lactoferrin as an active ingredient in producing a drug for improving lipid metabolism. to a patient in need thereof.

10. (Currently Amended) Use of lactoferrin as an-active-ingredient in producing a drug

A method for treating at least one disease or condition to be selected from the group consisting
of hypercholesterolemia, hyper-neutral lipidemia, hyper-low density lipoprotein (LDL)
cholesterolemia, hypo-high density lipoprotein (HDL) cholesterolemia, obesity, fatty liver and
cholesterol gallstone, said method comprising:

administering the composition of claim 2 having lactoferrin as an active ingredient to a patient in need thereof.

11. (Currently Amended) Use of lactoferrin as an active ingredient in producing a drug for treating a disease or condition for which A method for the improvement of basal metabolic rate, said method comprising; is to be effective.

administering the composition of claim 3 having lactoferrin as an active ingredient to a patient in need thereof.

12. (Currently Amended) The [[use]] method of any one of claims 9 to 11, wherein the composition is a drug is in the form of a dusting powder, a powder, a granule, a tablet or a capsule, and can be obtained by the steps comprising mixing the active ingredient with pharmaceutically acceptable additives in the dry state; if desired, subjecting the mixture to strong pressure molding in the dry state and successively forming the molded product into fine

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particulates or granules of a uniform-size; and tableting or encapsulating the mixture, the fine particulates or granules.

13. (Currently Amended) The use method of claim [[9,]] 12, wherein the composition is a drug is in the form of an enteric coated preparation.

14. (Canceled)

15. (Currently Amended) The use method of claim 9, wherein the drug is for the administration of the active ingredient is in an amount of about 0.1 mg to about 50,000 mg, preferably about 0.5 mg to about 10,000 mg, more preferably about 10 mg to about 2,000 mg a day.

16. (Canceled)

 (Withdrawn) A method of improving lipid metabolism comprising using lactoferrin as an active ingredient.

18. (Withdrawn) A method of treating at least one disease or condition to be selected from the group consisting of hypercholesterolemia, hyper-neutral lipidemia, hyper-low density lipoprotein (LDL) cholesterolemia, hypo-high density lipoprotein (HDL) cholesterolemia, obesity, fatty liver and cholesterol gallstone comprising using lactoferrin as an active ingredient.

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19. (Withdrawn) A method of treating a disease or condition for which the improvement

of basal metabolic rate is to be effective comprising using lactoferrin as an active ingredient.

20. (Withdrawn) The method of any one of claims 17 to 19, wherein the active

ingredient is used in the form of a dusting powder, a powder, a granule, a tablet or a capsule

which can be obtained by the steps of mixing the active ingredient with pharmaceutically

acceptable additives in the dry state; if desired, subjecting the mixture to strong pressure molding

in the dry state and successively forming the molded product into fine particulates or granules of

a uniform size; and tableting or encapsulating the mixture, the fine particulates or granules.

21. (Withdrawn) The method of claim 20, wherein the active ingredient is in the form of

an enteric coated preparation.

22. (Withdrawn) The method of claim 20, wherein the active ingredient is obtained by

coating tableted granules containing the active ingredient with a film having, as the main

component, a base which has resistance to the gastric juice and dissolves in the small intestine.

23. (Withdrawn) The method of claim 21 comprising administering the active ingredient

in an amount of about 0.1 mg to about 50,000 mg, preferably about 0.5 mg to about 10,000 mg,

more preferably about 10 mg to about 2,000 mg a day.

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24. (Withdrawn) The method of claim 22 comprising administering the active ingredient

in an amount of about 0.1 mg to about 50,000 mg, preferably about 0.5 mg to about 10,000 mg,

more preferably about 10 mg to about 2,000 mg a day.

25. (Withdrawn) The method of claim 21, wherein the active ingredient is in the form of

a dusting powder, a powder, a granule, a tablet or a capsule which can be obtained by the steps

comprising mixing the active ingredient with pharmaceutically acceptable additives in the dry

state; if desired, subjecting the mixture to strong pressure molding in the dry state and

successively forming the molded product into fine particulates or granules of a uniform size; and

tableting or encapsulating the mixture, the fine particulates or granules.

26. (Withdrawn) The method of claim 22, wherein the active ingredient is in the form of

a dusting powder, a powder, a granule, a tablet or a capsule which can be obtained by the steps

comprising mixing the active ingredient with pharmaceutically acceptable additives in the dry

state; if desired, subjecting the mixture to strong pressure molding in the dry state and

successively forming the molded product into fine particulates or granules of a uniform size; and

if desired, tableting or encapsulating the mixture, the fine particulates or granules.